

1 **Source differences in the components and cytotoxicity of PM<sub>2.5</sub> from**  
2 **automobile exhaust, coal combustion, and biomass burning**  
3 **contributing to urban aerosol toxicity**

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15 **Abstract.** Although air quality guidelines generally use the atmospheric concentration of fine particulate matter (PM<sub>2.5</sub>) as the  
16 metric for air pollution evaluation and management, the fact can't be ignored that different particle toxicity is unequal and  
17 significantly related to their sources and chemical compositions. Therefore, judging the most harmful source and identifying  
18 the toxic component would be helpful to optimize air quality standards and prioritize targeted PM<sub>2.5</sub> control strategies to protect  
19 public health more effectively. Since the combustions of fuels, including oil, coal, and biomass, are main anthropogenic sources  
20 of environmental PM<sub>2.5</sub>, their discrepant contributions to health risks of mixed ambient aerosol pollution dominated by  
21 respective emission intensity and unequal toxicity of chemical components need to be identified. In order to quantify the  
22 differences among these combustion primary emissions, ten types of PM<sub>2.5</sub> from each typical source group, i.e., vehicle exhaust,  
23 coal combustion, and plant biomass (domestic biofuel) burning, were collected for comparative study with toxicological  
24 mechanisms. Totally thirty type individual combustion samples were inter-compared with representative urban ambient air  
25 PM<sub>2.5</sub> samples, which chemical characteristics and biological effects were investigated by component analysis (carbon, metals,  
26 soluble ions) and *in vitro* toxicity assays (cell viability, oxidative stress, inflammatory responses) of human lung  
27 adenocarcinoma epithelial cells (A549). Carbonaceous fractions were plenteous in automobile exhaust and biomass burning,  
28 while heavy metals were more plentiful in PM<sub>2.5</sub> from coal combustion and automobile exhaust. The overall ranking of mass-  
29 normalized cytotoxicity for source-specific PM<sub>2.5</sub> was automobile exhaust > coal combustion > domestic plant biomass  
30 burning > ambient urban air, possibly with differential toxicity triggers, that the carbonaceous fractions (organic carbon, OC;  
31 elemental carbon, EC) and redox-active transition metals (V, Ni, Cr) assisted by water-soluble ions (Ca<sup>2+</sup>, Mg<sup>2+</sup>, F<sup>-</sup>, Cl<sup>-</sup>) might

32 play important roles in inducing cellular reactive organic species (ROS) production, causing oxidative stress and inflammation,  
33 resulting in cell injury and apoptosis, thus damage human health. Coupled with the source apportionment results of typical  
34 urban ambient air PM<sub>2.5</sub> in eastern China, reducing toxic PM<sub>2.5</sub> from these anthropogenic combustions will be greatly beneficial  
35 to public health. Besides the air pollution control measures that have been implemented, like strengthening the vehicle emission  
36 standards, energy switching from coal to gas and electricity, and controlling the open incineration of agricultural straws, further  
37 methods could be considered especially through preferentially reducing the diesel exhaust, then lessening the coal combustion  
38 by replacement with low-ash clean coals, and depressing the rural crop straw biomass burning emissions.

## 39 **1 Introduction**

40 As a mixture of multiple sources, ambient particulate matter (PM) arise from anthropogenic activities are continuously  
41 deteriorating the urban air quality, particularly in developing countries. Among these, fine PM with an aerodynamic diameter  
42 of less than 2.5  $\mu\text{m}$  (PM<sub>2.5</sub>) is recognized as a serious public health concern due to its long persistence in air, carcinogenicity  
43 and acute toxicity to humans (Al-Kindi et al., 2020). There were extensive epidemiological evidences that airborne PM can  
44 cause serious negative effects on human health, such as respiratory and cardiovascular diseases, genetic mutations, and  
45 developmental disorders (Chowdhury et al., 2022;Clemens et al., 2017;Lelieveld et al., 2021;Smith, 2021). Currently, either  
46 the world air quality guidelines or the national air quality standards use the mass concentration of PM<sub>2.5</sub> as the metric for PM<sub>2.5</sub>  
47 pollution evaluation and management, however, the particle toxicity are unequal and significantly related to their sources and  
48 chemical compositions varying with space and time (Shiraiwa et al., 2017). Therefore, to identify which component(s) and  
49 source(s) of ambient PM are most harmful to health, will be helpful to evaluate air quality and prioritize targeted PM control  
50 strategies for protecting public health more effectively.

51 Besides natural sources, most aerosols come from anthropogenic activities especially energy consumption, including the  
52 combustion of fossil fuels causing industrial emissions and automobile exhaust, and biomass burning (McDuffie et al.,  
53 2021;Wu et al., 2022). These diverse sources make the ambient air PM<sub>2.5</sub> become a complex mixture with multiple chemical  
54 components, such as salts, organic carbon (OC), elemental carbon (EC), mineral and trace metals (Bari and Kindzierski, 2016).  
55 The physiological mechanisms of PM-induced cell toxicity in respiratory system have been continuously investigated with  
56 some progresses (Li et al., 2022b;Kelly and Fussell, 2012, 2020;Shiraiwa et al., 2017;Mack et al., 2019), such as the metabolic  
57 activation, oxidative stress, inflammatory response, and apoptosis, focused on by current study. In brief, after inhalation and  
58 deposition onto the epithelium, redox-active materials in PM<sub>2.5</sub> can induce the release of reactive organic species (ROS), which  
59 cause oxidative stress (an imbalance between ROS and antioxidants, i.e., disequilibrium of the redox state of a cell) followed  
60 by inflammation and cell death. The ROS can mediate subsequent signaling pathways leading to biomolecule damage (e.g.,  
61 DNA, lipid, and protein) and cellular injury, through mediating inflammatory responses including the release of pro-  
62 inflammatory cytokines like IL-6 and TNF- $\alpha$  by epithelial cells (Ahmed et al., 2020;Landwehr et al., 2021). For instance,

63 oxidative stress could trigger the induction of pro-inflammatory transcription factors, such as nuclear factor (NF)- $\kappa$ B, via the  
64 mitogen-activated protein kinase (MAPK) signaling pathway. Components adsorbed on particle surface, such as redox-active  
65 metals (transition metals, Fe, Ni, V, Cr, Cu), organic compounds (polycyclic aromatic hydrocarbons, PAHs; quinones), or  
66 even carbonaceous core of particles, are responsible for oxidative stress (Ahmed et al., 2020; Cachon et al., 2014). The non-  
67 redox active metals (Zn, Pb, Al) can also influence the toxic effects of transition metals by exacerbating or lessening the  
68 production of free radicals. The EC may not be a directly toxic component of PM<sub>2.5</sub> but rather operate as a universal carrier of  
69 combustion-derived chemicals (semi-volatile organic fractions, transition metals) of varying toxicity (Kelly and Fussell, 2020).  
70 Inorganic soluble sulphates and nitrates are acidic and can interact with and influence the solubility other compositions like  
71 metal bioavailability (Fang et al., 2017; Weber et al., 2016). However, besides the well-known toxic pollutants in environment  
72 like heavy metals and PAHs, which specific components and which particular sources are the most critical factors dominating  
73 the ambient aerosols' health risks, still need be explored.

74 Past studies performed in various countries have focused on physicochemical characterization or biological effects of  
75 ambient air PM<sub>2.5</sub> respectively (Weagle et al., 2018; Jia et al., 2017; Wang et al., 2020). For example, the source analysis of  
76 PM<sub>2.5</sub> by photochemical modelling (Bao et al., 2018), chemical composition of regional PM<sub>2.5</sub> (Chi et al., 2022), and the  
77 mechanism of PM<sub>2.5</sub> toxicity was independently reported recently (Jia et al., 2020). Because differences in particle composition,  
78 sources, and toxicity appear in different urban environments (Zhao et al., 2019; Borlaza et al., 2018), the source profiles of  
79 different emission inventories were needed to elucidate the local aerosol pollution characteristics for control strategies. For  
80 instance, it was reported that increased hospital admission risks were significantly associated with sources of vehicle exhaust,  
81 coal combustion, and secondary inorganic aerosols; in particular, coal combustion was positively correlated with increases in  
82 mortality risks (Du et al., 2021). Coal combustion and vehicle exhaust contributed more significantly to cancer risks of  
83 respiratory exposure to atmospheric heavy metals in Tianjin city of the northern China during the cold seasons (31% and 11%)  
84 than the warm seasons (11% and 4%) (Tian et al., 2021); while in Nanjing city of the eastern China, traffic emissions and non-  
85 traffic combustion (coal/waste/biomass) contributed 35% and 31% to carcinogenic risks of urban PM<sub>2.5</sub>-associated metals  
86 respectively (Xie et al., 2020). Traffic was suggested playing the most crucial role in enhancing the toxicity of fine particles  
87 (Park et al., 2018). The particle composition of motor vehicle exhaust was related to automobile types with various fuels,  
88 engines, and loads (Lin et al., 2020). A strong catalytic reactivity of metals in PM emitted from diesel vehicles was observed  
89 by dithiothreitol (DTT) assay (Jesus et al., 2018). It was found that straw burning during the harvest season is a major trigger  
90 of severe air pollution in many regions (Sahu et al., 2021). Aerosols from open biomass burning in the Amazon had a stronger  
91 ability to induce ROS than laboratory-generated secondary organic aerosols (Tuet et al., 2019). Although there were emerging  
92 studies on particle emission from single source, quantitatively comparative studies on multi-source pollutants as well as the  
93 differential composition and unequal toxicity of various sources are still limited.

94 The main objective of current study was to compare the chemical components and corresponding mass-normalized  
95 toxicological effects of individual PM<sub>2.5</sub> from various combustion sources and their unequal contributions to ambient aerosol

96 health risks. The aim is to provide experimental evidences supporting the targeting control of specific anthropogenic sources  
97 with prominent risks based on their pivotal toxic components. Therefore, we collected both representative ambient PM<sub>2.5</sub>  
98 samples (n = 16) from urban air and typical source PM<sub>2.5</sub> samples (n = 30) from automobile exhaust, coal combustion, and  
99 plant biomass burning. Their independent profiles of chemical compositions and *in vitro* cytotoxicity (cell viability, oxidative  
100 stress, and inflammatory responses) were investigated and intercompared, to assess the differences in source-to-receptor  
101 toxicity and to infer the core toxic components and respective harmful contribution. The pivotal toxic components were  
102 identified based on the source-sink bi-directional composition-effect results, which were further used to assess the health  
103 toxicity contribution of various emission sources to ambient air PM<sub>2.5</sub>, supported by its source apportionment through positive  
104 matrix factorization (PMF) and chemical mass balance (CMB) models.

## 105 **2 Materials and methods**

### 106 **2.1 Collection of PM<sub>2.5</sub> samples from primary emissions of 30 typical combustion sources and from representative** 107 **ambient urban air**

108 Totally 30 types of primary PM<sub>2.5</sub> samples emitted directly from automobile exhaust, coal combustion, and plant biomass  
109 (domestic biofuel) burning were respectively collected as follows for both chemical and toxicological analyses.

110 A total of 10 types of vehicles were chosen for exhaust investigation. They were further categorized into 7 sub-groups,  
111 including small duty gasoline coaches (SDGCs), small duty diesel coaches (SDDCs), middle duty diesel coaches (MDDCs),  
112 heavy duty diesel coaches (HDDCs), light duty diesel vans (LDDVs), middle duty diesel vans (MDDVs), and heavy duty  
113 diesel vans (HDDVs). The detailed information of these representative local automobiles was showed in Table S1.

114 To cover all coal types consumed in the city, 10 representative types of coal were gathered for investigation. They were  
115 further classified into 4 sub-groups, including 2 types of honeycomb coal (HC), 3 types of anthracite coal (AC), and 2 types  
116 of bituminous coal (BC) mainly for restaurant or household use, and 3 types of industrial coal (IC) for coal-fired power plants  
117 and steel-smelting industry. The detailed characteristic to physical-chemical of these typical coals purchased from local market  
118 were showed in Table S2.

119 Considering the plant biomass combustion in rural areas surrounding the megacity, 10 representative types of agricultural  
120 and forestry solid wastes were gathered for investigation. Straws of rice, wheat, corn, soybean, peanut, rape, and sesame,  
121 corncob, branches of peach and pine, were selected as plant biomass fuels and further divided into 2 sub-groups, including 8  
122 types of crop straw and 2 types of firewood. The detailed characteristic analysis of these typical plant biomass fuels collected  
123 from rural areas around Nanjing city were showed in Table S3.

124 The PM<sub>2.5</sub> samples directly emitted from these combustion sources were collected by dilution channel sampling method  
125 (Figure S1), using a 4-channel particulate matter dilution sampler (HY-805, Hengyuan Technology Development Co., CN).  
126 Each sampling included 3 parallel channels of quartz microfiber filter (Figure S2) and 1 channel of Teflon membrane filter

127 with diameters of 47 mm, through a size selector for PM<sub>2.5</sub> with a flow rate of 160 L min<sup>-1</sup> (each channel is 40 L/min). Clean  
128 air was pumped for 10 min before and after each sample was collected. Before using, the blank quartz filters were incinerated  
129 by a muffle furnace at 500 °C for 3 h to remove any possible organic matters, while Teflon filters were baked at 60 °C for 4 h.  
130 After being equilibrated in a constant temperature and humidity chamber for 24 h, the filters were weighed both before and  
131 after sampling for gravimetric measurements, then the mass of collected PM<sub>2.5</sub> could be calculated. The sampled filters were  
132 stored in a refrigerator at -20 °C before analysis. The quartz filter loaded PM<sub>2.5</sub> samples were used for carbon and ion analysis,  
133 and for toxicity tests, while the parallel Teflon filter loaded samples were used to determine metals.

134 As the actual mixture of various source particles in real environment, totally 16 ambient air PM<sub>2.5</sub> samples (each time lasting  
135 23h) covering a year monthly were collected from December 2019 to October 2020 in an urban site surrounded by traffic,  
136 residential and commercial quarters of Nanjing city, Yangtze River Delta of eastern China, using a high-volume air sampler  
137 (800 L min<sup>-1</sup>) with quartz microfiber filters (Li et al., 2022a).

## 138 **2.2 Chemical composition analysis**

139 All collected source and ambient PM<sub>2.5</sub> samples were conducted following component analysis (Li et al., 2023). For the  
140 concentrations of heavy metals in particulates, samples were digested by concentrated HNO<sub>3</sub>-HClO<sub>4</sub> acids with a progressive  
141 heating program and determined by inductively coupled plasma optical emission spectrometry (ICP-OES; Optima8000,  
142 PerkinElmer, for Cr, Mn, Ni and Pb), with elements (V, Co, As) at lower concentrations measured by ICP mass spectrometry  
143 (ICP-MS; NexIONTM300X, PerkinElmer). Blank filter, reagent blank, replicates, and standard reference material (NIST SRM  
144 1648a, urban dust) were adopted for analytical quality control, with recoveries ranged 90-110 %. Carbonaceous species (OC  
145 and EC) in PM<sub>2.5</sub> were determined using a DRI-2001A OC/EC (Atmoslytic Inc., Calabasas, CA, USA). For the concentrations  
146 of water-soluble ions (WSIs), the main cations (Na<sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup>, NH<sub>4</sub><sup>+</sup>) and anions (NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>2-</sup>, Cl<sup>-</sup>, F<sup>-</sup>) in PM<sub>2.5</sub> were  
147 measured by ion chromatography (IC, Thermo Fisher Scientific, USA), using the Metrosep C6-150/4.0 column for cations and  
148 the Metrosep A Supp 5 150/4.0 column for anions, respectively.

## 149 **2.3 Preparing mass-normalized PM<sub>2.5</sub> suspension for cell exposure**

150 Totally 30 source and 16 ambient PM<sub>2.5</sub> samples were also performed cytotoxicity tests. In order to elute the particles  
151 completely from the quartz membranes, a whole PM<sub>2.5</sub>-loaded sample filter was cut into small pieces, immersed in ultrapure  
152 water and extracted six times (30 min for each) in an ultrasonic bath at 0 °C. Although the ultrasonication might impact the  
153 ROS (Miljevic et al., 2014), the inevitable systematical error was ignored in this study. The extract was then suction filtered  
154 through a 2.6 µm pore-size nylon membrane to remove possible quartz fragments, and the bulk filtrate was freeze-dried back  
155 to pure PM<sub>2.5</sub> powder. Ultimately, based on particle mass, the gathered PM<sub>2.5</sub> was dispersed by sterile phosphate-buffered  
156 saline (PBS) to a concentration of 400 mg L<sup>-1</sup>, and then diluted to PM<sub>2.5</sub> suspension of 80 mg L<sup>-1</sup> with serum-free Dulbecco's  
157 modified eagle medium (DMEM) for following *in vitro* cell exposure (Li et al., 2022a).

## 158 **2.4 Cell culture and cellular toxicity tests by *in vitro* PM<sub>2.5</sub> exposure**

159 Aerosol pollution can harm lung alveoli and epithelial cells, and the A549 adenocarcinoma epithelial cell has long been used  
160 as a suitable epithelial alveolar model (Li et al., 2022b; Park et al., 2018). The A549 cells were cultured in RMPI-1640 medium  
161 (Gibco, USA) supplemented with 10% fetal bovine serum (FBS, Hyclone, USA) and 1% antibiotic penicillin-streptomycin  
162 (100 U mL<sup>-1</sup>) at 37 °C in a 5% CO<sub>2</sub> incubator. After PM<sub>2.5</sub> exposure, cell viability and the indicators reflecting oxidative  
163 damage and inflammatory responses were determined respectively. While the cell viability assay was helpful in determining  
164 PM<sub>2.5</sub> dose to cells, the endogenous ROS measurements revealed the status of cellular oxidative potential after PM<sub>2.5</sub> exposure  
165 followed by the relative effects of ROS on various stages of cellular toxicity like inflammatory responses (Gali et al., 2019).  
166 The cell viability (metabolic activity) was evaluated by mitochondrial activity and determined by the methyl-thiazol-  
167 tetrazolium (MTT) assay (Chen et al., 2019). After trypsin action, the density of cells in the logarithmic growth phase was  
168 adjusted to 1 × 10<sup>5</sup> mL<sup>-1</sup>. Cell suspensions were inoculated into 96-well plates (Costar, USA) at 100 µL per well. The blank  
169 control well (without medium and PM<sub>2.5</sub> suspension) and reagent control well (with medium but without PM<sub>2.5</sub> suspension)  
170 were set together. After incubation for 24 h and removing the cellular supernatant, various types of PM<sub>2.5</sub> suspension  
171 (concentration of 80 mg L<sup>-1</sup>) were added to 96-well plates and incubated for 24 h. Based on pre-experiments, the oxidative  
172 stress and inflammation response sensitively under this dose, while the cell viability can keep sufficient. Fresh medium and  
173 MTT reagent (Solarbio, Beijing, CN) were added to each well and the supernatant was discarded, then 100 µL of formazan  
174 lysate was added to each well. The optical density (OD) values were measured at 490 nm using a microplate reader (Thermo  
175 MULTISKAN FC, USA). Cell viability (%) = (OD<sub>treatment</sub> - OD<sub>blank control</sub>) / (OD<sub>reagent control</sub> - OD<sub>blank control</sub>). The levels of cellular  
176 ROS production causing oxidative stress in cells, pro-inflammatory cytokines including tumor necrosis factor-alpha (TNF-α)  
177 and interleukin-6 (IL-6) production for determining the expression of genes related to the inflammatory response in the  
178 supernatant were analyzed by enzyme-linked immunosorbent assay (ELISA) kits (Jiangsu Meibiao Biotechnology Co., Ltd.,  
179 CN), and OD values were measured at 450 nm (Huang et al., 2020; Pang et al., 2020).

## 180 **2.5 Data analysis**

181 The statistical analysis was performed by IBM SPSS statistics 24 and plotted by Origin 2020b software. Spearman correlation  
182 coefficients were produced by the correlation analysis. The variance was statistically significant when the statistical test level  
183 was p < 0.05, and extremely significant when p < 0.01. Statistical analyses were performed using Kruskal-Wallis test (Kruskal  
184 and Wallis, 1952).

185 The source apportionment of PM<sub>2.5</sub> mass in urban ambient air was conducted by the receptor models PMF (EPA PMF  
186 version 5.0) and CMB (EPA CMB 8.0). All measured constituents (OC, EC, Cu, Cr, Co, Ni, As, Pb, Mn, V, Na<sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>,  
187 Ca<sup>2+</sup>, NH<sub>4</sub><sup>+</sup>, Cl<sup>-</sup>, F<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, and SO<sub>4</sub><sup>2-</sup>) were selected as PMF model input data, and a four-factor solution was chosen as the  
188 optimal solution based on an assessment of the interpretability of the source profiles and the seasonal variability of the source  
189 contributions. Due to the high concentration of sulfate and nitrate in ambient PM<sub>2.5</sub>, and being lack of specific actual source to

190 emit sulfate and nitrate, we added the virtual source profiles of secondary sources in CMB model (Table S4). The virtual source  
191 profiles of secondary sources are represented by the proportion of sulfate, nitrate and ammonium in pure ammonium sulfate  
192 and ammonium nitrate.

### 193 **3 Results**

#### 194 **3.1 Contributions of combustion primary sources to urban ambient air PM<sub>2.5</sub>**

195 As shown in Figure S3, although have been significantly improved with the national air quality in recent years, the estimated  
196 annual PM<sub>2.5</sub> concentrations of representative city Nanjing ( $59.1 \pm 20.5 \mu\text{g m}^{-3}$ ) was 1.7 times higher than the China national  
197 standard ( $35 \mu\text{g m}^{-3}$ ) and 11.8 times higher than the WHO guidelines ( $5 \mu\text{g m}^{-3}$ ). Urban air PM<sub>2.5</sub> pollution levels in the cold  
198 season were higher than the warm season. The similar source apportionment results from PMF and CMB models are illustrated  
199 in Figure 1. Four major sources of the ambient PM<sub>2.5</sub> were produced by the PMF model (Figure S4), including secondary  
200 aerosols, and primary particles of automobile exhaust, coal combustion, and plant biomass burning, which account for 34.0%,  
201 27.7%, 25.2%, and 13.1% of total PM<sub>2.5</sub> mass concentration, respectively. The CMB model source profiles are shown in the  
202 Table S4, and we normalized contribution of secondary aerosols (32.4%), and automobile exhaust (32.2%), coal combustion  
203 (25.1%), plant biomass burning (10.3%). Therefore, although the contribution of secondary aerosols cannot be ignored, the  
204 main anthropogenic sources of urban air PM<sub>2.5</sub> were primary emissions from the various fuel combustions.

#### 205 **3.2 Chemical compositions of different PM<sub>2.5</sub> from 30 combustion sources and from representative urban ambient air**

206 Typical chemical components including carbonaceous fractions, heavy metals, and WSIs of all PM<sub>2.5</sub> samples from both  
207 ambient air and combustion sources were analyzed and compared with each other.

208 According to the comparisons of PM<sub>2.5</sub> bound carbonaceous fractions (Figure 2), automobile and biomass sourced PM<sub>2.5</sub>  
209 contained significantly higher total carbon (TC) content than coal combustion and ambient air, while the OC/EC ratio trend  
210 was ambient air > coal combustion > biomass burning > automobile exhaust sources. It indicated that the carbon content of  
211 ambient PM<sub>2.5</sub> mixture was lower and dominated by OC than that of combustion primary sources, implying that the OC in  
212 ambient air may be aged or cleaned. The OC undergoes various chemical reactions in the atmosphere, such as oxidation by  
213 ozone and hydroxyl radicals, resulting in degradation. Figures S4-S7 showed the detailed carbon fraction characteristics  
214 (contents and ratio) of PM<sub>2.5</sub> from each specific source. Carbonaceous fractions in automobile exhaust PM<sub>2.5</sub> were high but the  
215 difference between OC and EC content was small. Depending on the diverse automobile fuels, loads and tailpipe emission  
216 standards, the concentrations of carbon fractions in exhaust PM<sub>2.5</sub> varied widely with vehicle categories. The carbonaceous  
217 portion of PM<sub>2.5</sub> gradually declines as emission regulations rise, and EC likewise declines dramatically (Figure S5). However,  
218 such differences among coal types were less, except the bituminous coal with extreme high OC (Figure S6). The carbonaceous

219 fraction of PM<sub>2.5</sub> from domestic plant biomass burning differed in raw material species that tree branches source PM<sub>2.5</sub>  
220 generally contained higher carbon contents than those from crop straws (Figure S7).

221 Based on the grouped (Figure 3) and individual (Figures S9-S12) distributions of the measured heavy metals in various  
222 PM<sub>2.5</sub>, the V concentrations of combustion sources were generally higher while Co and Mn were lower than ambient urban air.  
223 Coal combustion emissions carried highest levels of Pb and were enriched in Cu and As (Figure S10), while biomass burning  
224 were rich in Cr and Ni (Figure S11). However, automobile exhausts were enriched in most heavy metals, especially Cu, and  
225 Cr, Ni, V, Mn (Figure S9). Heavy metals from different types of automobile exhausts with the same emission standard varies  
226 greatly. Anthracite and industrial coal combustions contain similar heavy metals much more than bituminous coal. Generally,  
227 Pb, V, Mn, As, and Cu in branches source PM<sub>2.5</sub> were higher than straws, while Cr, Ni, and Co were dominant and higher in  
228 straw burning emissions. A special discovery was that corn cob burning PM<sub>2.5</sub> carried more heavy metals than corn straw and  
229 was the biomass with the highest emission levels of heavy metals. Correspondingly, ambient air PM<sub>2.5</sub> were also rich in most  
230 metals, especially Mn, Pb, and Ni, Cu, Cr. Therefore, coal combustion sources might contribute most Pb to urban ambient air,  
231 and contribute significant Cu and As with automobile exhaust emissions, while plant biomass burning and automobile sources  
232 contribute the Cr and Ni. Besides natural dust, automobile exhaust should be the main anthropogenic source of airborne Mn.  
233 Considering the PMF source apportionments of ambient aerosols, automobile exhaust should be the main source of Cr in urban  
234 air PM<sub>2.5</sub>, and also the source for Cu together with coal combustion.

235 According to the comparisons of water-soluble cation and anion concentrations in various PM<sub>2.5</sub> (Figure 4), coal  
236 combustions contained highest SO<sub>4</sub><sup>2-</sup> and NH<sub>4</sub><sup>+</sup>, automobile exhausts had highest contents of NO<sub>3</sub><sup>-</sup>, Na<sup>+</sup> and Ca<sup>2+</sup>, while plant  
237 biomass burning sources contained highest K<sup>+</sup> and Cl<sup>-</sup>, but Mg<sup>2+</sup> was the lowest for all sources. However, the urban ambient  
238 air PM<sub>2.5</sub> contained highest NO<sub>3</sub><sup>-</sup> and were also dominated by SO<sub>4</sub><sup>2-</sup> and NH<sub>4</sub><sup>+</sup>, for which NO<sub>3</sub><sup>-</sup> should be mainly contributed  
239 by secondary aerosols and automobile primary source, SO<sub>4</sub><sup>2-</sup> and NH<sub>4</sub><sup>+</sup> should be significantly from coal combustions. Besides  
240 NO<sub>3</sub><sup>-</sup>, Na<sup>+</sup> and Ca<sup>2+</sup>, automobile source PM<sub>2.5</sub> also had the highest F<sup>-</sup> and Mg<sup>2+</sup> concentrations than other sources. The detailed  
241 concentration distributions of WSIs in PM<sub>2.5</sub> from each specific source were provided in Figures S12-S14. The WSIs levels  
242 vary widely with specific source categories. PM<sub>2.5</sub> from LDDVs-2 had the lowest amount of WSIs compared to the other  
243 automobile exhausts (Figure S13). Similar to the metal composition, bituminous coal also had the lowest WSIs among all coals  
244 (Figure S14). Compared to branches, PM<sub>2.5</sub> from burning crop straws had much greater levels of K<sup>+</sup>, Cl<sup>-</sup>, SO<sub>4</sub><sup>2-</sup> and less levels  
245 of F<sup>-</sup>, NO<sub>3</sub><sup>-</sup> (Figure S15).

246 To summarize, the overall concentrations of measured TC, cumulated heavy metals and WSIs in PM<sub>2.5</sub> from each source  
247 type were showed in Figure 5. Among all source emission and environmental receptor samples, the cumulated heavy metals  
248 from coal combustion was highest and automobile exhaust was higher than ambient PM<sub>2.5</sub>, the overall carbon contents from  
249 automobile exhaust and biomass burning were both higher than ambient PM<sub>2.5</sub>, while only the cumulated soluble ions in PM<sub>2.5</sub>  
250 from primary source of coal combustion was equivalent to the ambient aerosols. In a word, chemical compositions of PM<sub>2.5</sub>  
251 distributed much diversely and varied significantly with the specific source types of combustion emissions.



### 252 **3.3 Cell viability, oxidative stress and inflammation levels exposed to various mass-normalized PM<sub>2.5</sub>**

253 Multiple toxicological endpoints (cell viability, oxidative stress, and inflammation) that facilitate identifying the specific  
254 particle triggering ROS and inflammatory responses resulting in cell death were evaluated for source-specific PM<sub>2.5</sub>. After 24  
255 h exposure to the same dose of different PM<sub>2.5</sub> obtained from specific emission sources, the A549 lung cells also showed varied  
256 toxicological responses (Figure 6). The survival rate of cells exposed to automobile exhaust PM<sub>2.5</sub> was much lower than  
257 ambient air PM<sub>2.5</sub> by 16.6% (Figure 6a). Automobile exhaust PM<sub>2.5</sub> induced the highest ROS production in cells higher than  
258 biomass burning, which was 26.4% and 14.8% higher than ambient PM<sub>2.5</sub> (Figure 6b). Coal combustion induced the highest  
259 cellular IL-6 production followed by automobile exhaust, which was 13.1% and 4.48% higher than ambient air PM<sub>2.5</sub>,  
260 respectively; while the PM<sub>2.5</sub> from automobile exhaust and biomass burning induced similarly 10.4% higher cellular production  
261 of TNF- $\alpha$  than ambient PM<sub>2.5</sub> (Figure 6c, 6d). These results suggested that, combustion primary emission PM<sub>2.5</sub> had stronger  
262 ability to induce oxidative stress and inflammatory injury in lung cells than ambient air PM<sub>2.5</sub>, thus resulted in the higher  
263 probability of apoptosis induction (Victor and Gottlieb, 2002; Wang et al., 2013). Generally, the mass-normalized PM<sub>2.5</sub> from  
264 primary source of automobile exhaust posed the strongest overall toxicity. Therefore, to protect public health by controlling  
265 PM<sub>2.5</sub> pollution, these anthropogenic combustions were key target sources, especially the most toxic automobile PM<sub>2.5</sub> should  
266 be reduced preferentially.

### 267 **3.4 Correlations between various PM<sub>2.5</sub> components and toxicity endpoints**

268 Spearman correlation coefficients between chemical compositions and cellular toxicological response indicators were applied  
269 to screen the key components of all PM<sub>2.5</sub> involved in cell injury (Figure 7). It was found that, the degrees of correlations  
270 varied with the toxicological mechanisms of different airborne chemicals. Based on the overall PM<sub>2.5</sub> samples from various  
271 sources, the pro-inflammatory cytokine IL-6 showed significantly strong positive correlations with some heavy metals (As,  
272 Pb, V, Cu), while TNF- $\alpha$  and oxidative stress (ROS) had similar significantly positive correlations with aerosol components  
273 of carbon fractions (EC, OC) and transition metals (V, Cr, Ni). The TNF- $\alpha$  also showed positive correlation with water soluble  
274 Cl<sup>-</sup> and K<sup>+</sup>, and ROS correlated with F<sup>-</sup>, Ca<sup>2+</sup> and Mg<sup>2+</sup>.

## 275 **4 Discussion**

### 276 **4.1 Chemical markers for source apportionments of ambient air PM<sub>2.5</sub>**

277 Combustion emissions are key anthropogenic sources contributing to urban air PM<sub>2.5</sub>, through both primary and secondary  
278 aerosols, which were 66% and 34% estimated by PMF model, 67.6% and 32.4% by CMB model, respectively (Figure 1).  
279 Compared to the PMF results, the proportions of coal combustion and secondary sources in the CMB results show minimal  
280 changes, while biomass contributions are slightly underestimated, and there is a slight increase in the proportion attributed to  
281 vehicular emissions. The high concentrations of chemical markers are usually used in source analysis, such as ammonium

282 sulfate and nitrate for secondary aerosols which are originated mainly from the gaseous precursors (e.g.,  $\text{NH}_3$ ,  $\text{SO}_2$  and  $\text{NO}_x$ )  
283 (Mahilang et al., 2021), the EC, Cu, Mn, and Ni for vehicle exhaust (Srivastava et al., 2021), the As, Pb, OC, EC,  $\text{SO}_4^{2-}$  and  
284 relatively low  $\text{NO}_3^-/\text{SO}_4^{2-}$  ratios for coal combustion (Dai et al., 2020), soluble  $\text{K}^+$  and  $\text{Cl}^-$  for plant burning (Jain et al., 2020).  
285 The detailed chemical species of these specific source emission  $\text{PM}_{2.5}$  samples also supported the results. Moreover, low  
286 OC/EC ratio of high TC content, high  $\text{NO}_3^-$ ,  $\text{F}^-$ ,  $\text{Na}^+$ ,  $\text{Ca}^{2+}$  and  $\text{Mg}^{2+}$ , V and Mn of automobile exhaust; Pb and As,  $\text{SO}_4^{2-}$  and  
287  $\text{NH}_4^+$  of coal combustion; soluble  $\text{K}^+$  and  $\text{Cl}^-$ , and high OC/EC ratio of high TC for plant biomass burning found in current  
288 study (Figures 2-5), could also be corresponding potential aerosol source markers.

#### 289 **4.2 Common $\text{PM}_{2.5}$ components related to specific combustion sources**

290 Generally, the automobile exhaust  $\text{PM}_{2.5}$  had high TC content and low OC/EC value with considerable EC content (Figure 2),  
291 varying with specific vehicle types (Figure S5-8). The contents of the carbon fractions from diesel vehicles were 2.39 times  
292 more than gasoline exhausts (Figure S5), and the OC/EC ratios of diesel exhausts were 37.3% of gasoline vehicles, owing to  
293 both considerable contents of EC and OC from diesel vehicle emission  $\text{PM}_{2.5}$ . Some diesel vehicles showed higher EC  
294 emissions with age, so exhaust cleaning devices for them are suggested. In addition, the amounts of OC and EC in exhausts  
295 gradually decreased with the strengthened emission standards they met (Wong et al., 2020). In  $\text{PM}_{2.5}$  samples obtained from  
296 coal combustion (Figure S6), the TC contents of bituminous coals was 3.97, 6.41, and 11.6 times higher than that of honeycomb  
297 coals, anthracite coals, and industrial coals, respectively, because bituminous coals contain higher volatile fraction. Emissions  
298 of non-methane VOCs increase with the volatile content of the coal (He et al., 2022). The vast majority of organic aerosols  
299 from bituminous coal are generated in the ignition and fierce combustion phases, which account for 99.9% of the entire  
300 combustion process; while these two phases of anthracite coal generate only 77% of the entire process (Zhou et al., 2016).  
301 Moreover, as the volatile matter in the coal decreases, the temperature at which weight loss begins and ends shift to higher  
302 values, that may be due to the lower amount of aliphatic chains present. It has been reported that for bituminous maximum  
303 weight loss happens in the range 490–600 °C, while in the case of anthracites coals it occurs between 750 and 870 °C (De la  
304 Puente et al., 1998). Therefore, besides the way of combustion and the use of combustion stoves, the coal quality related to  
305 different coal types and origins determine the carbonaceous fractions of the PM emitted by coal combustion (Zhang et al.,  
306 2022). In the  $\text{PM}_{2.5}$  samples from plant biomass combustion (Figure S7), OC contents were 2.21 times higher than EC contents,  
307 except that pine branches contained higher EC and rapeseed straw had considerable contents of EC and OC. The OC in ambient  
308  $\text{PM}_{2.5}$  dominated the carbonaceous component (Figure S8), consistent with the North China Plain and Indo-Ganges Plain  
309 (Flores et al., 2020; Xu et al., 2019). Combining the TC contents and OC/EC ratios, carbonaceous components in ambient  $\text{PM}_{2.5}$   
310 mainly originate from semi-volatile organic compounds (SVOCs) (Wang et al., 2018). Previous studies have reported that  
311 carbonaceous aerosols are mainly originated from fossil fuel combustion in transportation, coal combustion in power plants  
312 and industries, and biomass combustion (Kang et al., 2018; Zhang et al., 2015). Thus, to control ambient carbon aerosol

313 pollution, besides reducing the precursor emissions of secondary organic aerosols (SOA), controlling primary aerosols  
314 especially EC from diesel vehicles might be effective measures.

315 Airborne redox-active metals are usually linked with the oxidation stress of  $PM_{2.5}$ . Different types of automobiles emitted  
316 diverse metal contents (Figure S9). Metal elements in automobile exhaust are primarily contributed by fuels, lubricants, and  
317 engine component abrasion. Because Mn is a common antidetonator that delays and prevents the oxidation of hydrocarbons  
318 and increases the octane number, which not only increases the thermal efficiency of the engine but also improves the emission  
319 performance of the vehicle (Cheung et al., 2010), the Mn content was greater in gasoline vehicle exhausts than in diesel  
320 vehicles. Although there are multi-sources of traffic Pb emissions such as fuel combustion and brake wear (Wang et al.,  
321 2019;Panko et al., 2019), the automobile exhaust Pb content of gasoline vehicles were greater than diesel vehicles owing to  
322 oil combustion. Moreover, for the same vehicle type (LDDVs-1 and 2; HDDVs-1 and 2; SDGCs-1 and 2), the stricter the  
323 emission standard required, the lower the exhaust metal contents. The metal contents in the  $PM_{2.5}$  of trucks was higher than  
324 that of passenger cars (Wu et al., 2016). In the combustion  $PM_{2.5}$  of 10 coal types (Figure S10), Pb contents were the highest  
325 than other heavy metals, similar to available findings (Zhang et al., 2020). The  $PM_{2.5}$  metals from bituminous coal were  
326 significantly lower than other coal types, because indicated by the coal quality analysis, bituminous coal has a low ash content  
327 which is mainly derived from non-combustible minerals in coal. These findings suggested that coal maturity might be an  
328 important factor influencing the metal composition of particulates emitted from coal combustion (Shen et al., 2021;Zhang et  
329 al., 2021). Heavy metal contents in biomass burned  $PM_{2.5}$  varied much widely with raw plant types (Figure S11), although  
330 dominated by Cr and Ni. Different plant species and even different plant parts differ significantly in their ability to uptake and  
331 accumulate metals from soil (Zhao et al., 2020). Moreover, because of the high enrichment factors of some metals for crop  
332 straws (Zhang et al., 2016;Sun et al., 2019), they also released more Cr, Ni, and Co during burning than fuelwoods. Total metal  
333 emissions were highest in corn cob but lowest in peanut straw burning  $PM_{2.5}$ . The heavy metals enriched in urban ambient air  
334  $PM_{2.5}$  showed slightly seasonal pattern (Figure S12), while contents of V, Co, and As were relatively low and less affected by  
335 seasonal changes. Accordingly, supported by the metal profiles of anthropogenic combustion sources and ambient aerosols, to  
336 control the environmental airborne heavy metal pollution, key targets might be the Pb, Cu and As from honeycomb, anthracite  
337 and industrial coal combustion, Cu from vehicle exhausts and especially V from light duty diesel van with the CN.III emission  
338 standard and Mn from gasoline vehicles, Cr and Ni from biomass especially crop straws burning.

339 Epidemiological studies have also shown the mortality closely related to the WSIs such as sulfate and nitrate in aerosols  
340 (Ostro et al., 2009;Liang et al., 2022). Among the WSIs contents of various automobile exhaust  $PM_{2.5}$  (Figure S13),  $NO_3^-$  and  
341  $Ca^{2+}$  were the most abundant anion and cation, respectively. The high  $NO_3^-$  in the automobile  $PM_{2.5}$  may be due to  $NO_x$   
342 production during high-temperature combustion, while the high  $Ca^{2+}$  content should be related to additives in automobile fuels  
343 and calcium-based lubricants (Hao et al., 2019;Yang et al., 2019). Moreover, the exhaust WSIs decreased with the strengthened  
344 automobile emission standards required. Coal combustion  $PM_{2.5}$  contained relatively higher  $SO_4^{2-}$  and  $NH_4^+$  concentrations  
345 followed by  $Cl^-$  than other WSIs species (Figure S14). Among various coal types, industrial coals emitted highest  $SO_4^{2-}$

346 followed by honeycomb and industrial coal with also high  $\text{NH}_4^+$ , but bituminous coals emitted low WSIs which were mainly  
347  $\text{NO}_3^-$ ,  $\text{F}^-$  and  $\text{Na}^+$ ,  $\text{Ca}^{2+}$ . The WSIs emission factors of honeycomb coal were generally higher than those of lump coal (Yan et  
348 al., 2020). For biomass combustion emissions (Figure S15),  $\text{Cl}^-$  and  $\text{K}^+$  were dominant WSIs in  $\text{PM}_{2.5}$  from straw-type fuels  
349 (Tao et al., 2016; Sillapapiromsuk et al., 2013), but fuelwood-type combustion emitted high  $\text{NO}_3^-$ . Plant species absolutely  
350 determine the emissions (Liao et al., 2021). Finally, there were also high levels of  $\text{NO}_3^-$ ,  $\text{SO}_4^{2-}$ , and  $\text{NH}_4^+$  in ambient air  $\text{PM}_{2.5}$   
351 (Zhang et al., 2019) (Figure S16), even higher than the investigated combustion sources, so other sources like the secondary  
352 aerosols may also contribute. Consequently, target combustion primary aerosols WSIs might include, the  $\text{NO}_3^-$  from vehicle  
353 exhausts and fuelwood burning;  $\text{SO}_4^{2-}$  and  $\text{NH}_4^+$  from honeycomb, anthracite and industrial coal combustion;  $\text{Cl}^-$  and  $\text{K}^+$  from  
354 biomass especially crop straw burning.

### 355 **4.3 $\text{PM}_{2.5}$ toxicity related to specific sources by pivotal chemical components**

356 The complexity of the sources and compositions of atmospheric  $\text{PM}_{2.5}$  leads to different toxicological effects (Newman et al.,  
357 2020; Kelly, 2021). The toxicological effects of  $\text{PM}_{2.5}$  are not comparable among different studies owing to distinct exposure  
358 concentrations, biological models, endpoints, and  $\text{PM}_{2.5}$  generation methods (Kelly and Fussell, 2020; Park et al., 2018). In this  
359 study, we employed same exposure conditions and biological endpoints, in order to obtain comparable toxicity data for  $\text{PM}_{2.5}$   
360 from different sources. Our mass-normalized results demonstrated that automobile exhaust  $\text{PM}_{2.5}$  induced the highest lethality  
361 and cellular ROS and  $\text{TNF-}\alpha$  production, coal combustion  $\text{PM}_{2.5}$  induced the highest cellular IL-6 production, plant biomass  
362 burning  $\text{PM}_{2.5}$  induced considerable cellular  $\text{TNF-}\alpha$  and ROS production (Figure 6). Generally, various toxicities of  
363 combustion emission primary  $\text{PM}_{2.5}$  were much greater than the urban ambient air  $\text{PM}_{2.5}$  (Figure 6), owing to the higher  
364 concentrations of specific toxic components in  $\text{PM}_{2.5}$  from these sources. The supplementary information had included  
365 exhaustive cytotoxicity indicators from each individual source (Figure S17-S20). While the survival rate of cell exposed to  
366 CN.III emission standard  $\text{PM}_{2.5}$  was the lowest and the capacity to induce cells to produce ROS was the highest for CN.IV,  
367 automobile exhaust had a similar potential to cause cells to produce inflammatory cytokines (Figure S17). The capability to  
368 induce IL-6 production in cells was highest for industrial coal  $\text{PM}_{2.5}$ , whereas bituminous coal had the highest survival rate of  
369 cells and  $\text{TNF-}\alpha$  induction capacity (Figure S18). From the Figure S19 we can see that the  $\text{PM}_{2.5}$  cytotoxicity of straws and  
370 branches burning was analogous, but it should be noted that the cell viability of various straw  $\text{PM}_{2.5}$  differs significantly, that  
371 may be related to the raw fuel characteristics.

372 These possible mechanisms were implied by the overall relationships between the measured chemical components with  
373 cytotoxicity indicators of  $\text{PM}_{2.5}$  from various specific sources (Figure 7). In general, both  $\text{TNF-}\alpha$  and ROS were significantly  
374 positively correlated with carbonaceous fractions and redox-active transition metals (V, Cr, Ni), which were main contributors  
375 of automobile exhausts and biomass burning. The IL-6 was significantly positively correlated with some heavy metals (As and  
376 Pb, V and Cu), which were main contributors of coal combustion sources. Potential mechanisms include that, carbon fractions  
377 bound in  $\text{PM}_{2.5}$  could be transformed into reactive metabolites and then induce ROS production in cells (Stevanovic et al.,

378 2019), and the PM<sub>2.5</sub> bound transition metals could also induce ROS production through the Fenton reaction and disrupt the  
379 function of enzymes in cells (Verma et al., 2010; Sørensen et al., 2005). Oxidative stress can lead to inflammatory infiltration  
380 of neutrophils and stimulate immune cells to produce inflammatory cytokines, among which TNF- $\alpha$  and IL-6 play important  
381 roles in the inflammation development (Xu et al., 2020). Ultimately, excessive production of ROS leads to dysfunctional  
382 endoplasmic reticulum responses and dysfunctional lipid metabolism in ROS bursts can result in cell membrane damage and  
383 even cell death (Piao et al., 2018; Zhao et al., 2004). There have been some related supporting reports. For instance, the OC  
384 and EC were significantly associated with biological responses of PM from vehicle emissions collected in tunnels (Niu et al.,  
385 2020). The polar or quinone fractions of PAHs in diesel engine exhaust particles significantly contributed to the heightened  
386 toxic response (Xia et al., 2004). The PM<sub>2.5</sub> generated from biomass burning contained a substantial concentration of  
387 carbonaceous components. In addition, Cr and Ni in PM<sub>10</sub> from straws were highly associated with ROS (Li et al., 2023). In  
388 current study, cellular ROS was also correlated with water soluble Ca<sup>2+</sup>, F<sup>-</sup>, and Mg<sup>2+</sup>, which were main contributors of  
389 automobile exhaust PM<sub>2.5</sub>. The Ca<sup>2+</sup> controls the membrane potential and regulates mitochondrial adenosine triphosphate (ATP)  
390 production, and excessive Ca<sup>2+</sup> leads to energy loss and more ROS production (Madreiter-Sokolowski et al., 2020). Moreover,  
391 the TNF- $\alpha$  was also positively correlated with water soluble Cl<sup>-</sup> and K<sup>+</sup>, which were main contributors of plant burning PM<sub>2.5</sub>.  
392 Therefore, the accumulations of some organic matters with high carbonaceous content (OC, EC) in PM<sub>2.5</sub> typically from  
393 automobile exhausts and plant biomass burning, redox-active metals (V, Cr, Ni) and water-soluble anions (Cl<sup>-</sup>, F<sup>-</sup>) and cations  
394 (Ca<sup>2+</sup>, Mg<sup>2+</sup>) contributed by various combustions, might induce ROS production in cells, cause cellular damage through  
395 oxidative stress and inflammatory responses, impair cell viability and finally harm human health.

396 Considering the multi-endpoints measured and the PM<sub>2.5</sub> toxicity mechanisms mentioned above, based on the cell viability  
397 first, and then ROS followed by inflammatory markers, together with the significantly related toxic chemical composition  
398 contents (Park et al., 2018), we put forward a general sequence of overall mass-normalized toxicity for these combustion  
399 source PM<sub>2.5</sub> to managers. To improve the urban environmental air quality for better public health benefits by controlling  
400 aerosols pollution, considering the differential toxicity intensity of each chemical component and their contributions from  
401 various sources to ambient aerosols, preferential targets of specific primary PM<sub>2.5</sub> sources and bound pollutants from  
402 anthropogenic combustions are suggested as following sequence: reducing the automobile exhaust PM<sub>2.5</sub> containing high  
403 contents of EC, transition metals (V, Cu, Ni, Cr), and ions (Ca<sup>2+</sup>, Mg<sup>2+</sup>, F<sup>-</sup>, Na<sup>+</sup>) from diesel exhausts by strengthening the  
404 emission standards and accelerating the phasing out of highly polluting vehicles; then lessening the coal combustion rich in  
405 heavy metals (As, Pb, Cu) by replacement with low-ash clean coals; and depressing the biomass burning containing high OC,  
406 Ni, Cr, Cl<sup>-</sup> and K<sup>+</sup> from rural crop straw emissions and promoting domestic cleaner energy such as natural gas.

#### 407 **4.4 Limitations and perspectives**

408 In current study, we selected A549 cell based on previous abundant experimental experiences and also because it has been  
409 used popularly in *in vitro* toxicology studies to elucidate the cellular and molecular mechanisms of PM involved in lung for

410 many decades (Li et al., 2022b). However, recently the human normal bronchial epithelial cell BEAS-2B was preferred over  
411 the human lung adenocarcinoma epithelial cell A549. For instance, both cells were used in an aerosol study (Bonetta et al.,  
412 2017), results of which highlighted the higher sensitivity of BEAS-2B cells respect to A549 also in samples with low level of  
413 pollutants, because the PM<sub>0.5</sub> samples from Italian towns can induce genotoxicity in normal cells while cancer cells might be  
414 resistant to their adverse effects. Therefore, although our results are reasonable under the same exposure conditions, there were  
415 still potential limitations of A549 cells since they may be more resistant to exposure to external compounds, and the generally  
416 more sensitive BEAS-2B cells are suggested for future studies.

417 In toxicity assessments, cell vitality reflects the overall health of cells, encompassing factors such as cell membrane integrity,  
418 intracellular metabolic activity, and cell proliferation capacity. Decreased cellular vitality may be associated with cell damage,  
419 toxic effects, or cellular apoptosis. Inflammation markers are employed to assess the extent and nature of inflammatory  
420 reactions, including the production of cytokines and inflammatory mediators, as well as the activation status of inflammatory  
421 cells. Inflammation is a complex physiological response, typically delineated by the immune and inflammatory reactions of  
422 the body to stimuli such as injury or infection. Alterations in inflammation markers can indicate the intensity and nature of the  
423 inflammatory response. In this study, multiple biological responses of epithelial cells to various PM<sub>2.5</sub> were evaluated,  
424 including that, cell viability evaluated the mitochondrial dehydrogenase activity of the living cells, excessive intracellular ROS  
425 formation induced by PM<sub>2.5</sub> was responsible for oxidative stress to the cells, cytokines IL-6 and TNF- $\alpha$  were determined for  
426 the effect of PM<sub>2.5</sub> on pro-inflammatory response in cells. In general, in vitro data can be used to rank various types of particles  
427 in terms of the toxic potential including possible carcinogenicity. Each marker will help to understand the hazard and toxicity  
428 of PM<sub>2.5</sub>. However, the toxicity of PM<sub>2.5</sub> may be the result of multiple components acting through disparate physiological  
429 mechanisms, with inconsistent relationships among endpoints (Park et al., 2018). For instance, in BEAS-2B cells, oxidative  
430 stress generated by H<sub>2</sub>O<sub>2</sub> exposure often results in cytotoxicity rather than by stimulating cytokine/chemokine responses,  
431 sometimes no correlation between oxidative damage and cytokine/chemokine responses. Moreover, TNF- $\alpha$  gene was not  
432 detected in BEAS-2B cells exposed to atmospheric PM collected from Benin, but the gene expression of other inflammatory  
433 cytokines (IL-1 $\beta$ , IL-6, and IL-8) were significantly induced, and decreasing cell viability was highly correlated with high  
434 secretion of all studied cytokines (Cachon et al., 2014). Therefore, in the present study, it was impossible to analyze all  
435 chemicals in PM<sub>2.5</sub> and determine all related toxicological endpoints, so unmeasured chemicals and endpoints might also play  
436 roles in the incongruous or unexplained results, and we also can't over-explain the mechanisms just based on statistical  
437 relations. To overcome these hurdles, standardization of toxicological studies (experimental methodologies) and reporting  
438 guidelines are necessary for tracking and comparing results.

439 This study ranked the unequal "toxic effects" based on the same mass concentration of PM<sub>2.5</sub> exposure in body lung fluid  
440 system, while the "health risks" usually relating to the inhalation exposure concentration of PM<sub>2.5</sub> in ambient air were not  
441 calculated and evaluated quantitatively. Moreover, non-linear concentration-response functions for various endpoints and  
442 different exposure concentrations might also limit using toxicological data straightforwardly to predict health effects

443 (morbidity, mortality) in human populations, so drawing conclusions precisely quantifying/ranking the health risks of PM<sub>2.5</sub>  
444 from specific sources or of individual PM<sub>2.5</sub> components is still not an easy task (Kelly and Fussell, 2020). Therefore, coupled  
445 with source apportionment and exposure level of ambient aerosols pollution, toxicology combined with epidemiology studies  
446 linking these factors and indicating scientific mechanisms would help to reach conclusions.

447 Moreover, the exact effective measures to control these specific key toxic components from the emissions of various  
448 combustion sources indeed a challenge, but still need to be explored. The findings of this research provide a specific direction  
449 for better air pollution control and public health. Besides the environmental technological methods of controlling toxic  
450 components targeting source materials, combustion processes, and final emissions, the environmental management policies  
451 are also beneficial to such aims, like the choice of fuel types, especially for the management of domestic biomass fuel burning.  
452 For examples, potential solutions include promoting new green energy vehicles and low-ash clean coals, depressing the diesel  
453 exhaust and rural crop straw burning emissions.

## 454 **5 Conclusions**

455 In current study, we found that 2/3 mass of urban ambient air PM<sub>2.5</sub> in a typical megacity of eastern China originated from  
456 primary sources of anthropogenic combustions including coal, automobile, and biomass. Because of the significant differences  
457 in the chemical compositions, the diverse PM<sub>2.5</sub> from both mixed ambient air and directly from individual combustion sources  
458 showed much differential mass-normalized *in vitro* toxicity to the human lung epithelial cells, either for the environmental  
459 aerosol samples collected from different seasons, or for the primary emissions of PM<sub>2.5</sub> from various specific source types.  
460 According to the comparative study and correlation analysis, the carbonaceous fractions (OC, EC) and redox-active heavy  
461 metals (V, Ni, Cr) assisted by water-soluble ions (Ca<sup>2+</sup>, Mg<sup>2+</sup>, F<sup>-</sup>, Cl<sup>-</sup>) might play important roles in inducing cellular ROS  
462 production, causing oxidative stress and inflammation, resulting in cell injury and apoptosis, thus damage human health. These  
463 toxic pollutants accumulated in specific-source PM<sub>2.5</sub> varied by the emission types and raw fuel properties. Combined with  
464 chemical composition and general cytotoxicity rank, the preferential controlling targets of specific combustion sources might  
465 be automobile exhaust (diesel vehicles with emission standards inferior to CN.IV), coal combustion (high ash and high sulfur  
466 coals), and rural plant biomass burning (crop straws). Although showing the synthetic effects of mixed compositions and  
467 complex sources, besides preventing the secondary aerosols from combustions, preferentially targeted reductions of toxic  
468 PM<sub>2.5</sub> direct emissions from these primary sources, would produce great benefits for public health with improved ambient air  
469 quality. Overall, the chemical findings of our toxicological research could help to support the precise, oriented, effective,  
470 efficient, and economical composition-source-based strategies for urban aerosols pollution control. However, as a prospect,  
471 the detailed mechanisms for unequal toxicity of PM with complicated components from various sources and their quantitative  
472 contributions to the health effects of ambient air PM<sub>2.5</sub> mixture still need in-depth study.

473 **Supplementary materials**

474 There are 20 figures (Figure S1-S20) and 3 tables (Table S1-S4) in the Supporting Information.

475 **Data availability**

476 All raw data can be provided by the corresponding authors upon request.

477 **Author contributions**

478 XSL conceived and supervised the study; WH, YP, MT, HL, and ZZ collected the samples; WH, YP, MT, WL, HL, ZZ, GS,  
479 and LX analyzed the chemical compositions; WH, YP, and MT performed the toxicity tests; WH, YP, MT, and XSL analyzed  
480 the data; WH and XSL wrote the manuscript draft; XSL, WH, GS, and TM reviewed and edited the manuscript.

481 **Competing interests**

482 The authors declare that they have no conflict of interest.

483 **Financial support**

484 This work was supported by the National Natural Science Foundation of China (NSFC 41977349, 41471418).

485



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## 738 Captions of figures

739 **Figure 1.** Source contributions (%) to the urban ambient air PM<sub>2.5</sub> (models PMF vs CMB).

740 **Figure 2.** Carbon contents (mg kg<sup>-1</sup>) and ratio in PM<sub>2.5</sub> from various specific sources (n=10 for each combustion source and  
741 n=16 for urban ambient air).

742 **Figure 3.** Heavy metal contents (mg kg<sup>-1</sup>) in PM<sub>2.5</sub> from various specific sources (n=10 for each combustion source and n=16  
743 for urban ambient air).

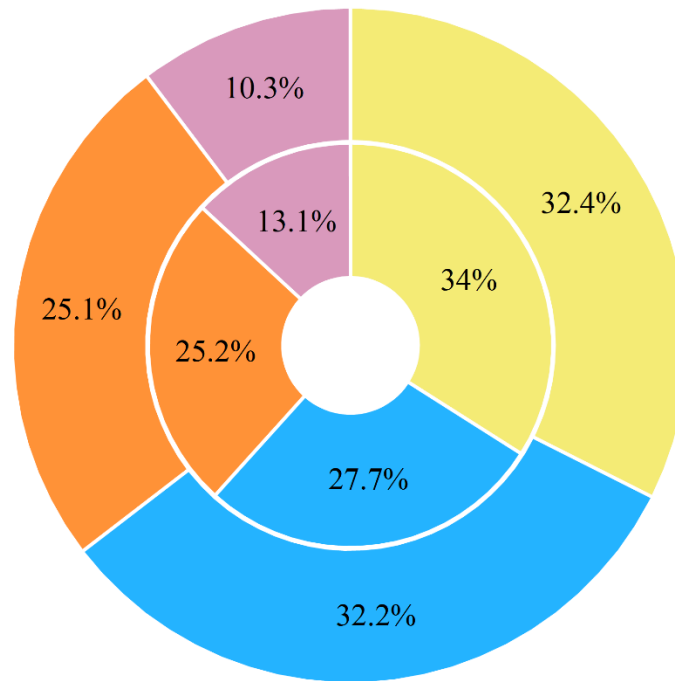
744 **Figure 4.** Water-soluble ion (WSI) contents (mg kg<sup>-1</sup>) in PM<sub>2.5</sub> from various specific sources (n=10 for each combustion  
745 source and n=16 for urban ambient air).

746 **Figure 5.** Cumulated typical measured components (mg kg<sup>-1</sup>) in PM<sub>2.5</sub> from various specific sources (n=10 for each  
747 combustion source and n=16 for urban ambient air).

748 **Figure 6.** Cell viability, oxidative stress and inflammation levels of human alveolar epithelial cell lines (A549) exposed to  
749 PM<sub>2.5</sub> suspension (80 mg L<sup>-1</sup>) from various specific sources (n=10 for each combustion source and n=16 for urban ambient  
750 air).

751 **Figure 7.** Overall correlations between typical cellular toxicological responses and chemical compositions of PM<sub>2.5</sub> from  
752 various sources (\*p < 0.05, #p < 0.01; n=46).

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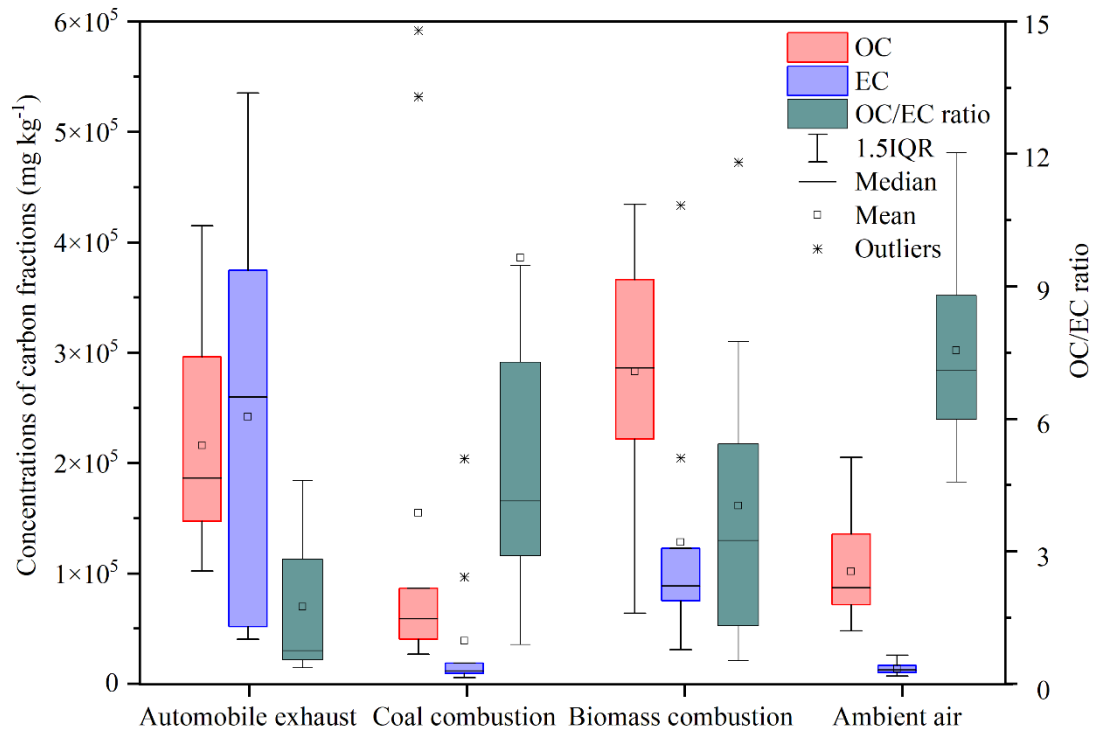
Inner ring - outer ring: PMF model -CMB model

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Secondary aerosols Automobile exhaust Coal combustion Biomass burning

756 **Figure 1: Source contributions (%) to the urban ambient air PM<sub>2.5</sub> (models PMF vs CMB).**

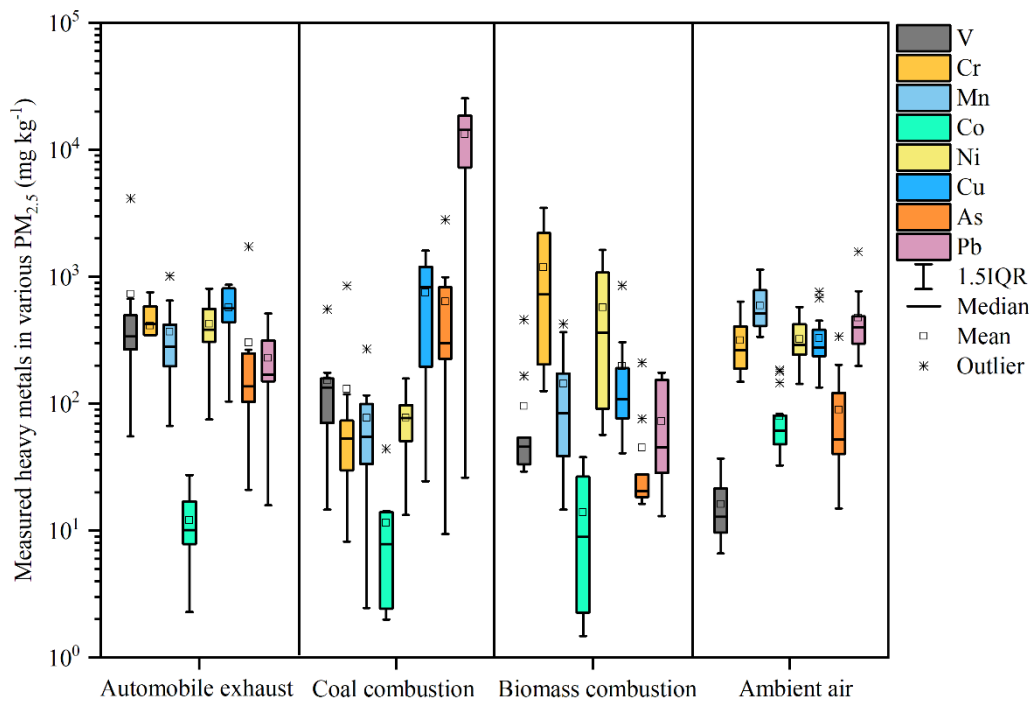
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760 **Figure 2: Carbon contents (mg kg<sup>-1</sup>) and ratio in PM<sub>2.5</sub> from various specific sources (n=10 for each combustion source and n=16**  
 761 **for urban ambient air).**

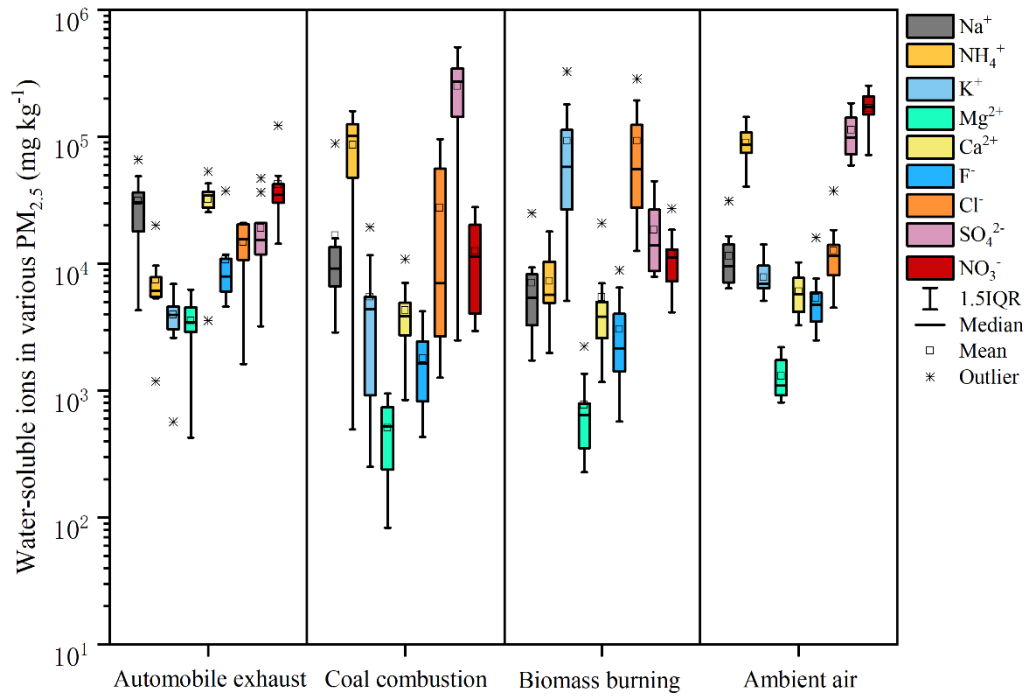
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764 **Figure 3: Heavy metal contents (mg kg<sup>-1</sup>) in PM<sub>2.5</sub> from various specific sources (n=10 for each combustion source and n=16 for**  
 765 **urban ambient air).**

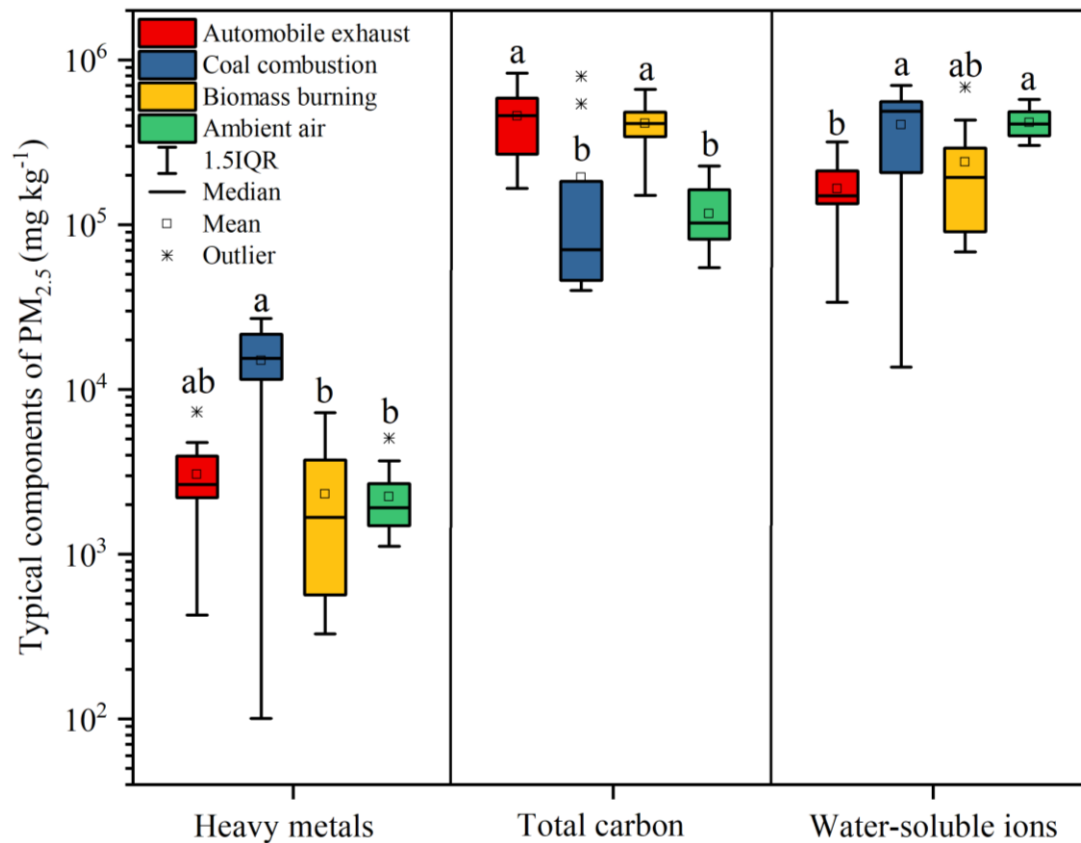
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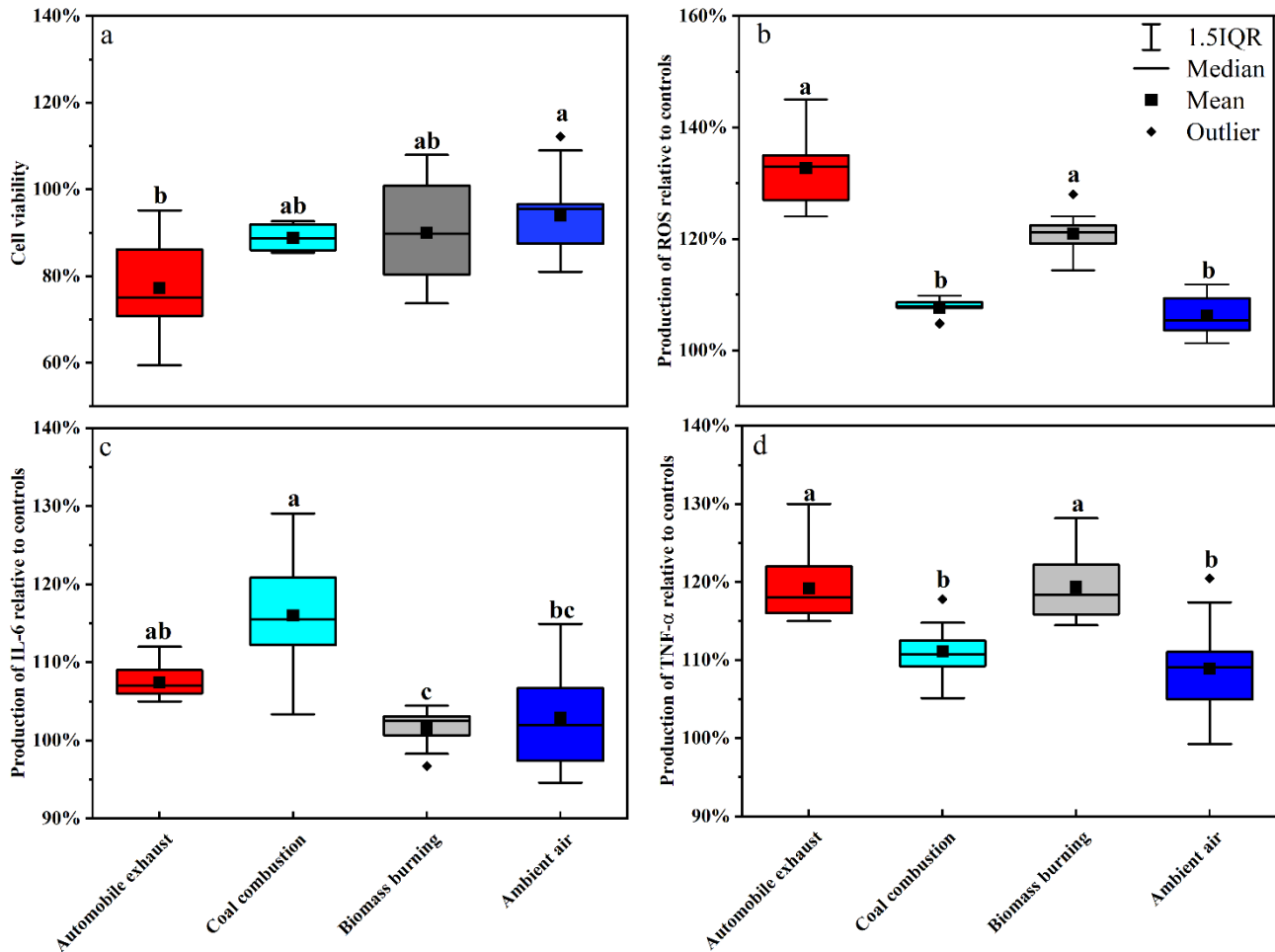
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772 **Figure 5: Cumulated typical measured components (mg kg<sup>-1</sup>) in PM<sub>2.5</sub> from various specific sources (n=10 for each combustion**  
 773 **source and n=16 for urban ambient air). Statistically significant differences between the groups are indicated by different letters**  
 774 **(Kruskal-Wallis test, p < 0.05).**

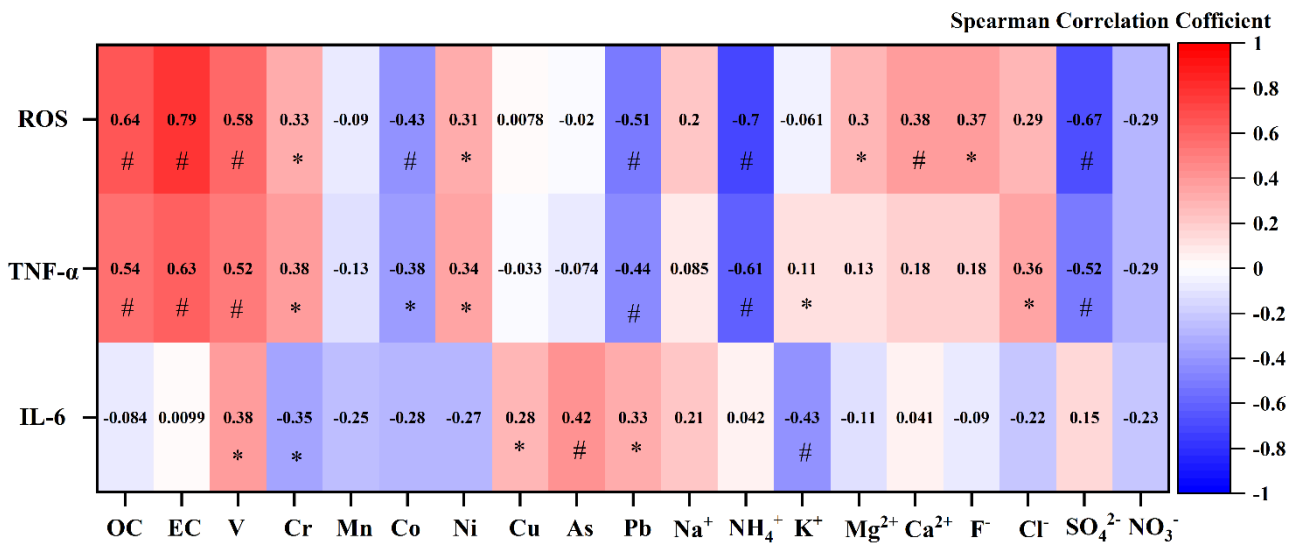


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777 **Figure 6: Cell viability, oxidative stress and inflammation levels of human alveolar epithelial cell lines (A549) exposed to PM<sub>2.5</sub>**  
 778 **suspension (80 mg L<sup>-1</sup>) from various specific sources (n=10 for each combustion source and n=16 for urban ambient air). Statistically**  
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